MQ - T PROPYL SILOXANE RESINS

CROSS REFERENCE TO RELATED APPLICATIONS [0001] Not applicable.

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FIELD OF THE INVENTION

[0002] The present invention provides siloxane resins, herein known as MQ-T propyl resins, comprising (R¹₃SiO_{1/2})_a, (R²₂SiO_{2/2})_b, (R³SiO_{3/2})_c, and (SiO_{4/2})_d units, where at least 40 mole % of the R³ groups are propyl. The present invention further relates to a method of preparing such siloxane resins by reacting a MQ siloxane resin with a T propyl siloxane resin. The siloxane resins of the present invention are useful in a variety of personal, household, automotive, and medical care applications, and in particular, as a resin additive in pigmented cosmetic formulations.

15 BACKGROUND OF THE INVENTION

[0003] Siloxane resins of the general formula R_nSiO_{(4-n)/2}, where R is an alkyl group and n is generally less than 1.8, are an important family of silicone polymers because of their utility in many commercial applications such as adhesive compositions and coatings applications. One particular subclass of siloxane resins, known as MQ resins (since they comprise primarily of "M" units of the general formula R₃SiO_{1/2} and "Q" units of the general formula SiO₂), have found utility in cosmetic formulations. In particular, MQ resins are commonly used in "extended wear" or "transfer resistant" cosmetic formulations. In these formulations, MQ resins enhance the substantivity of the pigments or other formulation actives to skin after application creating a longer lasting, and hence extended wear product.

[0004] Representative examples of transfer resistant cosmetic compositions using MQ resins are found in US 6,071,503, US 6,074,654, US 6,139,823, US 6,340,466, WO 97/17058, and WO 97/17059 which disclose compositions comprising the combination of organosiloxane resins and fluid diorganosiloxane resins with a volatile carrier.

[0005] US 5,330,747 teaches cosmetics with enhanced durability using a film forming agent from a pressure sensitive adhesive composition comprising a trimethylsilyl endblocked resinous copolymer, a silanol endblocked polydiorganosiloxane fluid, and a phenyl containing polysiloxane fluid.

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[0006] US 5, 075,103 and US 5,733,537 teach a hair treating method for imparting curl retention to hair in which at least one film forming ingredient is applied to the hair. The improvement utilizes as the film forming ingredient an organosilicon compound which is a nonpolar silsequioxane.

- 5 [0007] US 5,800,816 discloses cosmetic compositions having improved transfer resistance comprising: a) from about 0.1-60% by weight of trimethylated silica, b) from about 0.1-60% by weight of a volatile solvent having a viscosity of 0.5 to 100 centipoise (mPa·s) at 25°C, c) 0.1-60% by weight of a nonvolatile oil having a viscosity of 200 to 1,000,000 centipoise (mPa·s) at 25°C, d) 0.1-80% of a cosmetically acceptable carrier.
- 10 [0008] US 5,837,223 and US 6,036,947 teach transfer resistant high luster cosmetic stick compositions comprising, by weight of the total composition: a) 10-70% of a volatile solvent having a viscosity of 0.5 to 20 centipoise (mPa·s) at 25.degree. C., b) 0.5-40% of a guerbet ester, and c) 0.1-20% of a siloxysilicate polymer.
- [0009] GB 2,319,527 discloses fragrance releasing non-volatile polysiloxanes based on a high molecular weight polydiorganosiloxane compounds where at least one or more of the organic substituents of the polymer is a radical derived from a fragrant alcohol.
 - [0010] Japanese examined patent publication 1994-72085 teaches makeup cosmetic compositions having improved water resistance and durability containing an organic silicone resin, a volatile silicone oil, and a make up powder.
- 20 [0011] While the use of MQ resins in cosmetics have led to formulations having extended wear or transfer resistance, a need exists to alter the properties of the siloxane resins used in such formulations. In particular, films of MQ resins used in these formulations can have a matte finish and feel tacky. Thus, there is a need for improved siloxane resins that offer at least comparable extended wear and transfer resistance properties of the MQ resins presently used in cosmetic formulations, but having improved gloss (i.e. non-matte) that are non-tacky. Furthermore, there is a need for resins in hair care formulations that improve the curl retention properties of hair following treatment.
 - [0012] The present inventors have discovered improved siloxane resins by incorporating propyl containing siloxane (R³SiO_{3/2})_c units (T units) with M and Q units. The resulting siloxane resins, herein referred to as MQ T propyl siloxane resins, have improved physical properties. In particular, cosmetic formulations containing the present MQ-T propyl siloxane

resins have improved gloss vs MQ resins and are less tacky than T propyl resins, while maintaining their long lasting or wear characteristics.

SUMMARY OF THE INVENTION

- 5 [0013] This invention relates to a MQ-T propyl siloxane resin comprising the units:
 - (i) $(R^{1}_{3}SiO_{1/2})_{a}$
 - (ii) (R²₂SiO_{2/2})_b
 - (iii) $(R^3SiO_{3/2})_c$, and
 - (iv) $(SiO_{4/2})_d$
- 10 wherein

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 $R^1,\,R^2$, and R^3 are independently an alkyl group having from 1 to 8 carbon atoms , an aryl group, a carbinol group, or an amino group,

a has a value 0.05 to 0.5,

b has a value of zero to 0.3,

c has a value greater than zero,

d has a value of 0.05 to 0.6,

the value of a + b + c + d = 1,

with the proviso that greater than 40 mole% of the R³ groups in the siloxane resin are propyl. [0014] The invention further relates to a method of making a siloxane resin composition and the products obtained therefrom. The method comprises reacting:

A) a MQ resin comprising at least 80 mole % (R¹₃SiO_{1/2})_a and (SiO_{4/2})_d units where R¹ is an alkyl group having from 1 to 8 carbon atoms, an aryl group,

a carbinol group, or an amino group,

a and d has a value greater than zero,

the ratio of a/d is 0.5 to 1.5;

and

B) a T propyl resin comprising at least 80 mole % R³SiO units,

where R³ is an alkyl group having from 1 to 8 carbon atoms, an aryl group,

- a carbinol group, or an amino group,
- c has a value greater than zero,

and with the provisio that at least 40 mole % of the R³ groups are propyl,

WO 2005/075542 PCT/US2005/002451

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wherein the weight ratio of A/B is from 95:5 to 15:85.

[0015] The MQ-T propyl siloxane resins are useful in a variety of personal, household, or medical care compositions. In particular, the MQ-T propyl siloxane resins provide glossy, non-tacky films that can be used to enhance the substantivity of color cosmetic formulations.

The MQ-T propyl siloxane resins can also be used as additives in hair care formulations to enhance curl retention properties. The MQ-T propyl siloxane resins can also enhance the feel of treated hair, providing a softer feel compared to other siloxane resin additives. Thus, the present invention provides personal, household, or medical care compositions comprising the MQ-T propyl siloxane resins described herein.

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DETAILED DESCRIPTION OF THE INVENTION

[0016] The MQ-T propyl siloxane resin of the present invention comprises (i) (R¹₃SiO_{1/2})_a, (ii) (R²₂SiO_{2/2})_b, (iii) (R³SiO_{3/2})_c, and (iv) (SiO_{4/2})_d units which are commonly known in the art, and also used herein, as M, D, T, and Q units respectively. The amount of each unit present in the MQ-T propyl siloxane resin can be expressed as a mole fraction (i.e. a, b, c, or d) of the total number of moles of all M, D, T, and Q units present in the MQ-T propyl siloxane resin. The value of a (mole fraction of M units) is 0.05 – 0.5, or alternatively 0.15 to 0.4. The value of b (mole fraction of D units) is 0 – 0.3, alternatively 0 to 0.1, or alternatively 0 to 0.05. Thus, the MQ-T propyl siloxane resins can be free of D units, or alternatively can contain up to and including 0.3 mole fraction of D units. The value of c (mole fraction of T units) is greater than 0, alternatively 0.05 to 0.65, or alternatively 0.4 to 0.65. The value of d (mole fraction of Q units) is 0.05 to 0.6, alternatively 0.2 to 0.6, or alternatively 0.2 to 0.55.

[0017] The MQ-T propyl siloxane resins of the present invention are characterized by having at least 40 mole %, alternatively 50 mole %, or alternatively 90 mole % of the R³ alkyl groups on the T unit be propyl.

[0018] The R¹, R², and R³ in the units of the MQ-T propyl siloxane resin are independently an alkyl group having from 1 to 8 carbon atoms, an aryl group, a carbinol group, or an amino group. The alkyl groups are illustrated by methyl, ethyl, propyl, butyl, pentyl, hexyl, and octyl. The aryl groups are illustrated by phenyl, naphthyl, benzyl, tolyl, xylyl, xenyl, methylphenyl, 2-phenylethyl, 2-phenyl-2-methylethyl, chlorophenyl, bromophenyl and fluorophenyl with the aryl group typically being phenyl.

[0019] For the purposes of this invention a "carbinol group" is defined as any group containing at least one carbon-bonded hydroxyl (COH) radical. Thus the carbinol groups may contain more than one COH radical such as for example

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5 [0020] The carbinol group if free of aryl groups has at least 3 carbon atoms, or an arylcontaining carbinol group having at least 6 carbon atoms, The carbinol group free of aryl
groups having at least 3 carbon atoms is illustrated by groups having the formula R⁴OH
wherein R⁴ is a divalent hydrocarbon radical having at least 3 carbon atoms or divalent
hydrocarbonoxy radical having at least 3 carbon atoms. The group R⁴ is illustrated by
alkylene radicals such as -(CH₂)_x- where x has a value of 3 to 10,

[0021] The aryl-containing carbinol group having at least 6 carbon atoms is illustrated by groups having the formula R⁵OH wherein R⁵ is an arylene radical such as -(CH₂)_xC₆H₄- wherein x has a value of 0 to 10, -CH₂CH(CH₃)(CH₂)_xC₆H₄- wherein x has a value of 0 to 10, -(CH₂)_xC₆H₄(CH₂)_x- wherein x has a value of 1 to 10. The aryl-containing carbinol groups typically have from 6 to 14 atoms.

[0022] The amino group is illustrated by groups having the formula $-R^6NH_2$ or $-R^6NHR^7NH_2$ wherein R^6 is a divalent hydrocarbon radical having at least 2 carbon atoms and R^7 is a divalent hydrocarbon radical having at least 2 carbon atoms. The group R^6 is typically an alkylene radical having from 2 to 20 carbon atoms. R^6 is illustrated by ethylene, propylene, $-CH_2CHCH_3$ -, butylene, $-CH_2CH(CH_3)CH_2$ -, pentamethylene, hexamethylene, 3-ethyl-hexamethylene, octamethylene, and decamethylene.

[0023] R⁷ is typically an alkylene radical having from 2 to 20 carbon atoms. R⁷ is illustrated by ethylene, propylene, -CH₂CHCH₃-, butylene, -CH₂CH(CH₃)CH₂-, pentamethylene, hexamethylene, 3-ethyl-hexamethylene, octamethylene, and decamethylene.

- [0024] Typical amino groups are -CH2CH2CH2NH2 and
- -CH2(CH3)CHCH2(H)NCH3, -CH2CH2NHCH2CH2NH2, -CH2CH2NH2,
- -CH2CH2NHCH3, -CH2CH2CH2CH2NH2, -(CH2CH2NH)3H, and
- -CH2CH2NHCH2CH2NHC4H9.
- 5 [0025] Typically, R¹ is a methyl group, R² is a methyl or phenyl group, and R³ is a propyl group.
 - [0026] Any individual D, T or Q siloxane units of the MQ-T propyl siloxane resins can also contain a hydroxy group and/or alkoxy group. Such siloxane units containing hydroxy and/or alkoxy groups are commonly found in siloxane resins having the general formula R_nSiO_{(4-n)/2}.
- The hydroxy groups in these siloxane resins typically result from the reaction of the hydrolyzable group on the siloxane unit with water; the alkoxy groups result from incomplete hydrolysis when alkoxysilane precursors are used or from exchange of alcohol with hydrolysable groups. Typically, the weight percent of the total hydroxy groups present in the MQ-T propyl siloxane resin is 3 %, alternatively, 2 %, or alternatively, 1.5%. Typically, the weight percent of the total alkoxy groups present in the MQ-T propyl siloxane resin is up to 20 %, alternatively up to 10 %.
 - [0027] The molecular weights of the MQ-T propyl siloxane resins are not restricted, but typically the number average molecular weight (M_N) ranges from 3,000 to 10,000, or alternatively from 5,000 to 8,000.
- 20 [0028] The MQ-T propyl siloxane resins of the present invention can be prepared by any of the methods known in the art for preparing siloxane resins having the general formula $R_n SiO_{(4-n)/2}$ where R is an alkyl group and n is generally less than 1.8. Alternatively, the MQ-T propyl resins can be prepared by the methods described infra.
 - [0029] The MQ-T propyl resins of this invention are illustrated by:
- 25 MO-T propyl resins comprising the units;

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((CH_3)_3SiO_{1/2})_a,

(R^3SiO_{3/2})_c, where R^3 = CH_3CH_2CH_2-, and

(SiO_{4/2})_d
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MQ-T propyl resins comprising the units;

30 $((CH_3)_3SiO_{1/2})_a$, $((CH_3)_2SiO_{2/2})_b$,

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(R^3SiO_{3/2})_c, where R^3 = CH_3CH_2CH_2-, and
                (SiO<sub>4/2</sub>)<sub>d</sub>
       MQ-T propyl resins comprising the units;
                ((CH_3)_3SiO_{1/2})_a,
                ((CH_3)_2SiO_{2/2})_b, ((CH_3)(C_6H_5)SiO_{2/2})_{b'},
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                (R^3SiO_{3/2})_c, where R^3 = CH_3CH_2CH_2-, and
                (SiO_{4/2})_d
       MQ-T propyl resins comprising the units;
                ((CH_3)_3SiO_{1/2})_a,
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                ((CH_3)_2SiO_{2/2})_b,
                (R^3SiO_{3/2})_c, where R^3 = CH_3CH_2CH_2, and (C_6H_5SiO_{3/2})_c
                (SiO_{4/2})_d
       MQ-T propyl resins comprising the units;
                ((CH_3)_3SiO_{1/2})_a,
                ((CH_3)_2SiO_{2/2})_b, ((CH_3)(C_6H_5)SiO_{2/2})_{b'},
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                (R^3SiO_{3/2})_c, where R^3 = CH_3CH_2CH_2-, (C_6H_5SiO_{3/2})_c, and
                (SiO<sub>4/2</sub>)<sub>d</sub>
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wherein a has a total value in the resin of 0.05 to 0.5, the sum of b + b' has a total value in the resin of zero to 0.3, c has a total value in the resin of 0.05 to 0.65, and d has a total value in the resin of 0.05 to 0.6.

[0030] The present invention also provides a method of making a siloxane resin composition and the products obtained therefrom. The method comprises reacting:

A) a MO resin comprising at least 80 mole % (R¹₃SiO_{1/2})_a and (SiO_{4/2})_d units where R¹ is an alkyl group having from 1 to 8 carbon atoms, an aryl group,

a carbinol group, or an amino group, a and d has a value greater than zero,

the ratio of a/d is 0.5 to 1.5;

and

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B) a T propyl resin comprising at least 80 mole % R³SiO units, where R³ is an alkyl group having from 1 to 8 carbon atoms, an aryl group, a carbinol group, or an amino group, c has a value greater than zero,

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and with the provisio that at least 40 mole % of the R³ groups are propyl, wherein the weight ratio of A/B is from 95:5 to 15:85.

[0031] Component A) is a MQ resin comprising at least 80 mole % (R¹₃SiO_{1/2})_a and (SiO_{4/2})_d units where R¹ is the same as defined above, i.e. an alkyl group having from 1 to 8 carbon atoms an aryl group, a carbinol group, or an amino group, a and d has a value greater than zero, and the ratio of a/d is 0.5 to 1.5. MQ resins suitable for use as component (A), and methods for their preparation, are known in the art. For example, U.S. Patent No. 2,814,601 to Currie et al., November 26, 1957, which is hereby incorporated by reference, discloses that MQ resins can be prepared by converting a water-soluble silicate into a silicic acid monomer or silicic acid oligomer using an acid. When adequate polymerization has been achieved, the resin is end-capped with trimethylchlorosilane to yield the MQ resin. Another method for preparing MQ resins is disclosed in U.S. Patent No. 2,857,356 to Goodwin, October 21, 1958, which is hereby incorporated by reference. Goodwin discloses a method for the preparation of an MQ resin by the cohydrolysis of a mixture of an alkyl silicate and a hydrolyzable trialkylsilane organopolysiloxane with water.

[0032] The MQ resins suitable as component A) in the present invention may contain D and T units, providing that at least 80 mole %, alternatively 90 mole % of the total siloxane units are M and Q units. The MQ resins may also contain hydroxy groups. Typically, the MQ resins have a total weight % hydroxy content of 2-10 weight %, alternatively 2-5 weight %. The MQ resins can also be further "capped" wherein residual hydroxy groups are reacted further with M groups.

[0033] Component B) is a T propyl resin comprising at least 80 mole % of R³SiO_{3/2} units, where R³ is the same as defined above, i.e. an alkyl group having from 1 to 8 carbon atoms an aryl group, a carbinol group, or an amino group, and with the proviso that at least 40 mole % of the R³ groups are propyl. Typically, the T propyl resin is a silsesquioxane resin. Silsesquioxane resins are well known in the art and are typically prepared by hydrolyzing an organosilane having three hydrolyzable groups, such as a halogen or alkoxy group, present in the molecule. Thus, component (B) can be obtained by hydrolyzing propyltrimethoxysilane, propyltriethoxysilane, propyltripropoxysilane, or by co-hydrolyzing the aforementioned propylalkoxysilanes with various alkoxysilanes. Examples of these alkoxysilanes include methyltrimethoxysilane, methyltriethoxysilane, methyltriisopropoxysilane, dimethyldimethoxysilane, and phenyltrimethoxysilane. Propyltrichlorosilane can also be

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hydrolyzed alone, or in the presence of alcohol. In this case, co-hydrolyzation can be carried out by adding methyltrichlorosilane, dimethyldichlorosilane, phenyltrichlorosilane, or similar chlorosilanes and methyltrimethoxysilane, methyltriethoxysilane, methyltriisopropoxysilane, or similar methylalkoxysilane. Alcohols suitable for these purposes include methanol, ethanol, n-propyl alcohol, isopropyl alcohol, butanol, methoxy ethanol, ethoxy ethanol, or similar alcohols. Examples of hydrocarbon-type solvents which can also be concurrently used include toluene, xylene, or similar aromatic hydrocarbons; hexane, heptane, isooctane, or similar linear or partially branched saturated hydrocarbons; and cyclohexane, or similar aliphatic hydrocarbons.

10 [0034] The T-propyl resins suitable as component B) in the present invention may contain M, D, and Q units, providing that at least 80 mole %, alternatively 90 mole % of the total siloxane units are T units. The T-propyl resins may also contain hydroxy groups. Typically, the T-propyl resins have a total weight % hydroxy content of 3-8 weight %.

[0035] A polyorganosiloxane can optionally be added to the method of the present invention as component C). Polyorganosiloxane useful as component C) in the present invention comprise R²₂SiO_{2/2}, (i.e. D units) or R³SiO_{3/2} (T units). The polyorganosiloxane can be added to introduce various D and T units into MQ-T propyl resins to alter the properties of the resulting resins. The structure or formula of the polyorganosiloxane is not restrictive, providing the polyorganosiloxane contains some measurable quantity of R²₂SiO_{2/2}, units (i.e. D units) or R3SiO3/2 (T units), and the total amount of polyorganosiloxane added to the reaction of A) and B) does not provide more than 50 mole % D or T units into the reaction mixture. The polyorganosiloxane can contain any combination of M, D, T and Q units, providing at least some D or T units are present. Thus, the polyorganosiloxane can be selected from any of the fluid, gum, or resinous silicones known in the art containing D or T units, or combinations thereof. The D units typically contain methyl or phenyl as the R² substituents, which can be designated as DMe and DPh respectively, or any combinations thereof. The T units typically contain methyl or phenyl as the R3 substituents, which can be designated as T^{Me} and T^{Ph} respectively, or any combinations thereof. The polyorganosiloxane can be a linear polydiorganosiloxane fluid having a viscosity of 10 -1000 cS (mm²/s). Typically the polydiorganosiloxane fluid is polydimethylsiloxane, or

alternatively a polymethylphenylsiloxane. The polyorganosiloxane can also be an

WO 2005/075542 PCT/US2005/002451

organosilsequioxane resin. The organosilsequioxane resin typically is a methylsilsequioxane resin or a phenylsilsequixone resin.

[0036] Components A), B), and optionally C) can be reacted by any method in the art known to effect reaction of M, D, T, and Q siloxane units. Typically however, components A), B), and optionally C) are reacted by a condensation reaction in the presence of a catalyst. Typically the MQ resin is contained in an aromatic hydrocarbon or siloxane solvent. Suitable condensation reaction catalysts are base catalysts including metal hydroxides such as potassium hydroxide and sodium hydroxide; metal salts such as silanolates, carboxylates, and carbonates; ammonia; amines; and titanates such as tetrabutyl titanates; and combinations thereof. Typically the reaction of components A), B), and optionally C) is effected by heating the reaction mixture to temperatures ranging from 50 to 140°C, alternatively 100 to 140°C. The reaction can be conducted in a batch, semi-continuous, or continuous process. [0037] The weight ratio of component A) to component B) (i.e. A/B) in the reaction can vary from 95:5 to 15:85, alternatively 95:5 to 20:80, or alternatively 90:10 to 20:80. The amount of component C) can vary, but the amount of component C) added should introduce less than 30 mole% of additional D or T units, based on the total moles of siloxane units in the reaction mixture.

[0038] The MQ-T propyl resins of the present invention are useful in a variety of personal, household, automotive, or medical care applications. In particular, the MQ-T propyl siloxane resins provide glossy, non-tacky films that can be used to enhance the substantivity of color cosmetic formulations. The MQ-T propyl siloxane resins can also be used in hair care formulations to enhance curl retention properties. Thus, the present invention provides personal, household, automotive, or medical care compositions comprising the MQ-T propyl siloxane resins described herein.

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EXAMPLES

[0039] The following examples are presented to further illustrate the compositions and methods of this invention, but are not to be construed as limiting the invention. All parts and percentages in the examples are on a weight basis and all measurements were obtained at about 23°C, unless indicated to the contrary.

Materials

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- [0040] MQ Resin = a MQ resin having the formula $M_{0.43}Q_{0.57}$ and M_n = 3230 dissolved in xylene at 70.8 wt % solids. The MQ resin was prepared according to techniques taught by Daudt in US 2,676,182.
- 15 [0041] T propyl resin = propyl silsesquioxane resin at 74.8 wt % in toluene. The propyl silsesquioxane resin was prepared from the hydrolysis of propyl trichlorosilane.
 - [0042] Phenyl silsesquioxane resin = a phenyl silsesquioxane solid flake resin at 100 wt % solids prepared from the hydrolysis of phenyl trichlorosilane.

Example 1 Preparation of MQT^{Pr} Resins

Stark trap fitted with a condenser on top, was loaded with MQ Resin, T propyl resin, xylene and 1MKOH in water at the ratios as shown in Table 1. The Dean Stark trap was pre-loaded with xylene to insure a 50% solids level in the reaction flask was maintained. The mixture in the flask was maintained at reflux temperature (100-140°C) for at least 3 hours. Any water that formed in the reaction mixture was continuously removed as needed and trapped as an azeotrope in the Dean Stark trap. After 3 hours of reflux, water was removed from the trap, and heating continued for an additional 30 min. After allowing the mixture to cool, excess acetic acid was added to neutralize the KOH in the mixture. The mixture was then filtered to remove the salts formed by passing it through a pressure filter. A solvent exchange was completed by heating the mixture in a rotary evaporator under vacuum. After the majority of

xylene was removed, decamethylcyclopentasiloxane was added while continuing to remove any residual aromatic solvent. The structures of the resulting siloxane resins were characterized by ²⁹Si NMR spectroscopy and GPC and the results summarized in Table 4 below.

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Table 1

Example #	Wt Ratio MQ/T ^{Pr} resins added	Wt% MQ Resin	Wt% T Propyl Resin	Wt% Xylene	Wt% 1 <i>M</i> KOH	Wt% Acetic Acid
1-a	(85:15)	59.4	10.5	29.1	0.9	0.2
1- b	(50:50)	34.9	34.8	29.1	0.9	0.2
1- c	(30:70)	20.9	48.8	29.2	0.9	0.2
1-d	(95:5)	67.1	3.5	28.3	0.9	0.2
1- e	(100:0)	69.3	0	28.8	0.9	0.2

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Example 2 Preparation of $MQT^{Pr}T^{Ph}$ resins

[0044] A series of MQT^{Pr}T^{Ph} resins were prepared following the experimental procedure described in Example 1 using the formulations shown in Table 2. In this series, a phenyl silsesquioxane resin was added to incorporate additional phenyl containing T units into the siloxane resin. The structures of the resulting siloxane resins were characterized by ²⁹Si NMR spectroscopy and GPC, as summarized in Table 4 below.

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Table 2

Example #	Wt Ratio MQ/T ^{Pr} /T ^{Ph} resins	Wt% MQ Resin	Wt% T Propyl Resin	Wt% T Phenyl Resin	Wt% of Xylene	Wt of 1M KOH solution in water	Wt of Glacial Acetic Acid
2-a	(85:7.5:7.5)	59.4	5.2	3.7	30.6	0.9	0.2
2-b	(50:25:25)	34.7	17.4	12.4	34.4	0.9	0.2
2-c	(15:42.5:42.5)	10.5	29.6	21.0	37.8	0.9	0.2

Example 3 Preparation of MQT^{Pr}D resins

[0045] Two MQT^{Pr}D resins were prepared following the experimental procedure described in Example 1 using the formulations shown in Table 3. In this series, a 100 cst polydimethylsiloxane fluid was added to incorporate D units (Me₂SiO) into the siloxane resin. The structures of the resulting siloxane resins were characterized by ²⁹Si NMR spectroscopy and GPC, as summarized in Table 4 below.

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Table 3

Example #	Wt Ratio MQ/T ^{Pr} /D added	Wt% MQ Resin	Wt% T Propyl Resin	Wt% D	Wt% of Xylene	Wt of 1M KOH solution in water	Wt of Glacial Acetic Acid
3-a	(38:57:5)	26.9	39.7	2.5	29.9	0.9	0.2
3-b	(28.5:66.5:5)	14.1	32.9	2.5	30.0	0.9	0.2

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Table 4

Example #	Resin structure according to NMR Characterization	wt% OH	Mn	Mw	Mw/Mn
MQ resin	M ^{0.43} Q ^{0.57}		3230	1516	4.7 .
T propyl resin	T ^{P7} 1.0	7.0	3470	11400	3.3
l-a	M _{0.374} Q _{0.529} :T ^{PT} _{0.097}	1.4	5880	271000	46.1
1-b	M _{0.248} Q _{0.341} :T ^{Pr} _{0.412}	2.1	6640	3860000	581.3
1-c	$M_{0.162}Q_{0.217}:T^{PT}_{0.621}$	1.5	7600	25300000	3329
1-d	M 0.419 Q0.5485; TPT 0.03	1.5			
1-e	MQ	1.7	5200	28900	5.6
2-a	M _{0.396} Q _{0.523} :T ^{Pr} _{0.0491} :T ^{Ph} _{0.0339}	1.6	5200	26900	5.2
2-b	M _{0,272} Q _{0,347} :T ^{P7} _{0,220} :T ^{Ph} _{0,161}	0.8	5867	1640000	279
2-c	M 0.0000 118: TPT 0.457: TPh 0.334	0.8			
3-a	MQ:T ^{rr} :D ^{me2}		5455	303000	55.5
3-b	M _{0.155} Q _{0,201} :T ^{Pr} _{0.591} :D ^{Mc2} _{0.053}	1.8	5140	295000	57.4

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[0046] Films resulting from the siloxane resins of Examples 1-3 were evaluated for gloss and tack after coating from 35% solids in volatile solvent onto a Leneta chart, evaluated in a foundation (color cosmetic) for durability, and in a hair fixative for curl retention and conditioning. The results are summarized in Table 6. The siloxane resins were applied to the hair tresses as 6 wt. % solutions in decamethylcyclopentasiloxane or isododecane. Descriptions of the specific test methods and composition of the cosmetic formulation used in these evaluations follows Table 6.

10 Table 6

Example #	60° Gloss	Tack	Foundation	Curl	Hair	Hair
•	ŀ		Durability -	Retention -	Conditioning	Conditioning -
	1		ΔE (Change	Hair – % curl	- Wet Comb	Dry Comb
	l		in color)	maintained	% Reduction	% Reduction
			İ	after 5 hours	in combing	in combing
				at 95% RH	force	force
No Resin -	55 blank		9.2	30.5 -36.8	-95 (exposed	-50 (exposed
neg control	chart				to water	to water only)
J					only)	
MQ resin	34.8	Not tacky	3.8	53-58	-55.7	-1650
T propyl resin	78.4	Very tacky	11.0	44		<u></u>
1-a	72.6	Not Tacky	5.1	45.5		
1-b	74.5	slight, less	4.7	50.6	-65	-1648
		w/time				
1-c	75.2	slight, less	5.4	51.2	2.2	-1508
		w/time				
l-d	68.2	Not tacky				
1-e	68.7	Not tacky				
2-a	62.3	Not Tacky	3.1	41.3		
2-b	75.0	Not Tacky	4.0	33.2		
2-с	80.9	Not Tacky	6.9	33.9		
3-a	69.5	Not tacky	3.9			L
3-b	76.6	Not Tacky		46.9	-31.5	-337

Gloss Measurement

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- 1) Coat Leneta charts (Form N2C) with solution using a #8 Meyer rod
- 2) Allow chart to dry for 1 hour. Measure 60° gloss using a portable gloss meter at 3 points on the left 1/3 of the chart. Calculate the average of the 3 gloss values. Evaluate coating for tack, greasiness, fingerprint mark and if and how the coating rubs off the chart.
 - 3) 4 hours after the drawdown was completed, measure 60° gloss using a portable gloss meter at 3 points on the middle 1/3 of the chart. Calculate the average of the 3 gloss values. Evaluate coating for tack, greasiness, fingerprint mark and if and how the coating rubs off the chart.
 - 4) 24 hours after the drawdown was completed, measure 60° gloss using a portable gloss meter at 3 points on the right 1/3 of the chart. Calculate the average of the 3 gloss values. Evaluate coating for tack, greasiness, fingerprint mark and if and how the coating rubs off the chart.
- 15 5) Using the average gloss at the 3 different times, calculate the overall average.

Curl Retention Test Method

Materials

- Prepared natural virgin brown hair tresses or oriental hair of 2 g, 25 cm.
 - Comb Trade Mark: Ace; reference 2618/6 GB.
 - Humidity chamber to regulate temperature and humidity during test.

Procedure for pre-treating the swatches (washing):

- 25 1) Wet 5 tresses for 30 sec with tap water at 37°C.
 - 2) Lather the 5 tresses for 30 sec. With 5 g of the 30% SLS solution (Empicol LX28/Albright & Wilson), stroking the tresses downward, ensure you repeat the same movement for all the hair washed. Leave on hair for 30 sec.
 - 3) Rinse the tresses for 1 minute with tap water at 37°C.
- 30 4) Remove the excess of water by running the tresses between the two fingers 3 times.
 - 5) Allow the tresses to dry overnight on a paper towel at room temperature.

Procedure for treating the swatches with the resin and curling:

Blank or negative control is the solvent used in the treatments.

- 35 1) Dip 1 tress at a time 3 times in 37 °C tap water and remove excess of water by stroking the tress between 2 fingers
 - 2) Lay the tress down on a clean support and apply 100 microliters of a 6% resins solution all along the using a calibrated micropipette
 - 3) Detangle the tress completely
- 40 4) Roll the tress on rod spiral curler,
 - 5) Leave the swatches to dry overnight in an oven at 40 °C.

Test - curl retention measurements:

- 1. Start the humidity chamber 2 hours before the test set at 70% humidity and 25 C.
- 2. Carefully remove the roller from hair by twisting it slightly, 10 minutes before the start of the test. Cut the ends of the tress in order to make it even (cut as few as possible).
- 3. Ensure each tress is correctly curled.
- 4. Hang the tresses in the humidity chamber: the bottom of the wax sealing on the tress should be on the line "0" of the millimeter paper sheet in the back of the chamber.
- 5. Measure hair tress length at predetermined intervals of time, the length is measured as the distance between the bottom of the wax sealing and the bottom of the tress be aware that the bottom of the tress is going down, so the view angle is always be perpendicular to the glass.
 - 6. After the 5 hours, remove the tresses from the humidity chamber and measure the tress length at its maximum, by unrolling it completely. Calculate curl retention as described below.

Percent curl retention is calculated as follows:

max length - length at T = x% Curl Retention = X 100 max length - length at T = 0

Instron Test for Hair Conditioning for Dry or Wet Hair

25 Materials

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- Testing was performed on slightly bleached European.
- Comb Trade Mark: Unbreakable or Ace Brand (Fine tooth spacing: 16 teeth/25 mm.)
 Wide tooth spacing: 11 teeth/25 mm.)

30 Hair swatches preparation and Baseline testing:

- 1) Weigh out between 2.35 and 2.50 g. hair
- 2) Rough up lower half of plastic tab with sandpaper
- 3) Cut $\sim 1/2$ " off root end of hair.
- 4) Put glue in middle of sanded end of tab and lay hair on the glue. Squeeze more glue on top of hair; then using another plastic tab press glue into hair while evenly spreading the hair and glue onto the tab keeping ~ 1/8 to 1/4 inch away from sides.
 - 5) Cut tress to $6" \pm 0.25"$ from bottom of tab.
 - 6) Let glue dry overnight.
 - 7) Punch hole in middle of tab $\sim 1/4$ " from the top.
- 40 8) Wash all tresses according to the following procedure:
 - a) Wet hair tress under 40 degree C tap water for 15 seconds.
 - b) Using a pipette, apply 1.0 g of 9% Sodium Lauryl Sulfate (active) and stroke through tress for 30 seconds.
 - c) Rinse tress under the running water for 30 seconds.
- d) Place tresses on paper towel covered tray and dry overnight.

- e) Comb through tress three times with narrow teeth.
- 9) Measure force it takes to comb the untreated tresses before treatment using an Inston machine. Take force measurements on dry and wet hair as follows before treatment.

Ensure that the temperature and humidity inside the test room are appropriate for testing (approximately 70 deg F and 30% RH).

a) A 5 kilogram load cell is used for the testing

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- b) Place 2 combs in the holders with wide-spaced teeth on the left and fine teeth on the right. Secure combs with holder screws so that approximately 1" of combs are sticking out of the front of the holder and over-lapping each other. For untreated tresses, use the same combs for combing all tresses; but for treated tresses, use different combs for every different treatment.
- c) Dry Combing Procedure: Detangle hair by combing tress 3 times, Retangle hair by swirling tress clockwise 3 times and counter clockwise 3 times, place tress on hanger and use Instron to comb through tresses.
- d) Wet Combing Procedure: Wet hair by dipping in distilled water and detangle hair by combing tress 3 times, retangle hair by dipping in distilled water three times, remove excess water by passing tress through index and middle fingers twice, place tress on hanger and use Instron to comb tress.

20 Procedure for treating and testing the swatches with resin solution:

- 1) Wet the tresses for 15 sec under tap water at 40°C.
- 2) Remove the excess water by pulling tress through index and middle fingers.
- 3) Lay the tresses on a clean support and apply 100 microliters of 6% active silicone product using a calibrated micropipette.
- 25 4) Allow solution to dry on hair 60 minutes for ethanol-based solutions, overnight for aqueous to test on dry hair.
 - 5) Comb through tresses once before performing Instron study on both dry and wet hair see 9 from previous section.

Foundation Formulation

Pigment Premix:

50 wt% DC 245 Fluid

- 5 13.16 wt% Carde AS Titanium dioxide (caprylyl silane treated)
 - 11.41 wt% Carde AS Red Iron Oxide(caprylyl silane treated)
 - 18.26 wt% Carde AS Yellow Iron Oxide(caprylyl silane treated)
 - 7.17 wt% Carde AS Black Iron Oxide(caprylyl silane treated)

10 Procedure:

- 1) Place DC 245 fluid in Waring Blender
- 2) Add titanium dioxide and mix by pressing the pulse button for 2 seconds for 15 seconds total.
- 3) Add red iron oxide and mix the same as titanium dioxide
- 15 4) Continue with the other pigments
 - 5) When all materials have been dispersed, mix on high and shred for 30 sec to grind the pigments
 - 6) Place premix into a round glass jar and place on pail roller for 6 hours.

20 Phase A

20.50 wt% Pigment Premix

7.50 wt% DC 5225C

8 wt% of a 50% resin solids in solvent

25 Phase B

54.80 wt % DI Water

1.0 wt% NaCl

0.20 wt% Polysorbate 20

Procedure for Liquid Water in oil Foundation

- 1) Put pigment dispersion on roller for 1 hour.
- 2) Weigh out resin and solvent to make a 50% solids dilution. Use oven and wheel to mix
- 5 3) Combine ingredients in Phase A, mix until uniform using a dual blade, turbulent style mixing action.
 - 4) Combine ingredients in Phase B in separate beaker, mix until uniform using a magnetic stirrer
 - 5) Increase mixing speed of Phase A to 1376 rpm and very slowly add Phase B through an addition funnel. This addition should take 10 mins.

Continue mixing for an additional 10 min.

Foundation Durability Method: Gardner Abrasion Tester

- 1. Cut collagen into 3.5" x 3" pieces, place one on each of the 3" x 2.5" polycarbonate blocks and put in the humidity chamber overnight. This chamber must be at a constant 98% relative humidity level.
 - 2. Remove collagen and block from chamber. Secure collagen to block with Scotch tape taking care not to place any tape on the top of the block's surface.
- 3. Add approximately 1 gram of foundation to the collagen, beading it across the top of the block. Using a #8 Meyer rod, coat the collagen with the foundation by placing the rod on the bead of foundation and spreading it downward to the bottom of the block. The final coating weight should be approximately 0.2 grams. This operation may need to be repeated to obtain the proper coating weight. Remove any material from the sides of the block.
- 4. Allow sample on collagen to dry. Drying times vary with different samples. Entire sample must be free from any wetness before testing. Measure color of sample on collagen for the initial baseline color using a spectrophometer or colorimeter. L*, a*, and b* designate the place of the colored object in a tri-dimensional space.
 - 5. Place block with collagen face-up on the Gardner Abrasion Tester making sure that the block is in the tester. The soft side of Velcro is attached to the insult block to abrade or insult the foundation sample on the collagen. The insult block rubs back and forth across the foundation sample. One insult consists of one back and forth motion. Insult the foundation sample on the collagen 20 times. The machine can be stopped at certain intervals to measure the color.
- 35 6. After the foundation sample is insulted 20 times, the color is read as L*, a*, b* and the change in color, ΔE, is calculated (see equation below). The number of insults, coating weight, and repetitions can be changed to fit the needs of the material being tested. This is up to the discretion of the operator.

 ΔL^* , Δa^* and Δb^* = value after abrasion – value at initial baseline before abrasion. $\Delta E = (\Delta L^2 + \Delta a^{*2} + \Delta b^{*2})^{1/2}$

With larger ΔE 's, more foundation was removed and therefore the foundation is less durable.

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CLAIMS

1. A siloxane resin comprising the units:

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- (i) $(R^1_3SiO_{1/2})_a$
- (ii) $(R^2_2SiO_{2/2})_b$
- (iii) (R³SiO_{3/2})_c, and
- (iv) (SiO_{4/2})_d

wherein

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 $R^1,\,R^2$, and R^3 are independently an alkyl group having from 1 to 8 carbon atoms, an aryl group, a carbinol group, or an amino group,

a has a value 0.05 to 0.5,

b has a value of zero to 0.3,

c has a value greater than zero,

d has a value of 0.05 to 0.6,

the value of a + b + c + d = 1,

with the proviso that greater than 40 mole % of the R³ groups in the siloxane resin are propyl.

- 2. The siloxane resin of claim 1 wherein the siloxane resin is selected from
- 20 MQ-T propyl resins comprising the units;

 $((CH_3)_3SiO_{1/2})_a$,

 $(R^3SiO_{3/2})_c$, where $R^3 = CH_3CH_2CH_2$ -, and

(SiO_{4/2})_d

MQ-T propyl resins comprising the units;

25 $((CH_3)_3SiO_{1/2})_a$,

 $((CH_3)_2SiO_{2/2})_b$,

 $(R^3SiO_{3/2})_c$, where $R^3 = CH_3CH_2CH_2$ -, and

(SiO_{4/2})_d

MQ-T propyl resins comprising the units;

30 $((CH_3)_3SiO_{1/2})_a$,

 $((CH_3)_2SiO_{2/2})_b$, $((CH_3)(C_6H_5)SiO_{2/2})_{b'}$,

 $(R^3SiO_{3/2})_c$, where $R^3 = CH_3CH_2CH_2$, and

(SiO_{4/2})_d

MQ-T propyl resins comprising the units;

 $((CH_3)_3SiO_{1/2})_a$,

 $((CH_3)_2SiO_{2/2})_b$,

 $(R^3SiO_{3/2})_c$, where $R^3 = CH_3CH_2CH_2$ -, and $(C_6H_5SiO_{3/2})_c$

 $(SiO_{4/2})_{d}$

MQ-T propyl resins comprising the units;

 $((CH_3)_3SiO_{1/2})_a$,

 $((CH_3)_2SiO_{2/2})_b$, $((CH_3)(C_6H_5)SiO_{2/2})_{b'}$,

 $(R^3 {\rm SiO_{3/2}})_c$, where $R^3 = C H_3 C H_2 C H_2$ -, $(C_6 H_5 {\rm SiO_{3/2}})_c$, and

 $(SiO_{4/2})_{d}$

wherein a has a total value in the resin of 0.05 to 0.5, the sum of b + b' has a total value in the resin of zero to 0.3, c has a total value in the resin of 0.05 to 0.65, and d has a total value in the resin of 0.05 to 0.6.

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- 3. A method of making a siloxane resin comprising reacting:
 - A) a MQ resin comprising at least 80 mole % (R¹₃SiO_{1/2})_a and (SiO_{4/2})_d units where R¹ is an alkyl group having from 1 to 8 carbon atoms, an aryl group, a carbinol group, or an amino group,

a and d has a value greater than zero, and

the ratio of a/d is 0.5 to 1.5;

and

B) a T propyl resin comprising at least 80 mole % R³SiO units,

where R3 is an alkyl group having from 1 to 8 carbon atoms,

an aryl group, a carbinol group, or an amino group,

c has a value greater than zero,

and with the provisio that at least 40 mole % of the R³ groups are propyl, wherein the weight ratio of A/B is from 95:5 to 15:85.

- 30 4. A siloxane resin prepared by the method of claim 3.
 - 5. A personal care product comprising the siloxane resin of claim 1 or 4.

- 6. The personal care product of claim 5, where the personal care product is a cosmetic product.
- 7. The personal care product of claim 5, where the personal care product is a hair care product.

INTERNATIONAL SEARCH REPORT

Intermional Application No
PCT/US2005/002451

A. CLASSIF IPC 7	COSG77/44 COSG77/04 A61K7/11	C08G77/10						
	International Patent Classification (IPC) or to both national classification	on and IPC						
	B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols)							
IPC 7								
Documentat	ion searched other than minimum documentation to the extent that su	ch documents are included in the fields sea	arched					
Electronic da	ata base consulted during the international search (name of data base	e and, where practical, search terms used)						
EPO-In	ternal, WPI Data, PAJ							
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT							
Category °	Citation of document, with indication, where appropriate, of the rele	vant passages	Relevant to claim No.					
A	EP 0 336 633 A (LOCTITE CORPORATION) 1-7 11 October 1989 (1989-10-11)							
A	example 1 US 2002/031488 A1 (KANJI MOHAMED ET AL) 14 March 2002 (2002-03-14) paragraph '0016!; example 1							
Furt	ther documents are listed in the continuation of box C.	Patent family members are listed in	in annex.					
"A" docum consis "E" earlier filing "L" docum which citatic "O" docum	*Special categories of cited documents: *A' document defining the general state of the art which is not considered to be of particular relevance *E' earlier document but published on or after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone which is cited to establish the publication date of another citation or other special reason (as specified) *O' document reterring to an oral disclosure, use, exhibition or other means *P' document published prior to the international filing date but later than the priority date and not in conflict with the application but cited to understand the principle or theory underlying the considered novel or cannot be considered to involve an inventive step when the document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined with one or more other such document is combined to involve an inventive step when the document is combined to involve an inventive step when the document is combined to involve an inventive step when the document is combined to involve an inventive step when the document is combined to involve an inventive							
Date of the	actual completion of the International search	Date of mailing of the international sea	arch report					
3	3 June 2005	14/06/2005						
Name and	Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (-31-70) 340-3016 Buestrich, R							

INTERNATIONAL SEARCH REPORT

rormation on patent family members

PCT/US2005/002451

Patent document cited in search report		Publication date		Patent family member(s)		Publication date
EP 0336633	A	11-10-1989	AT	141305	T	15-08-1996
EI 0330030	••		DE	68926925	D1	19-09-1996
			DE	68926925	T2	06-02-1997
			ĒΡ	0336633	A2	11-10-1989
			ËS	2090034	T3	16-10-1996
			HK	1005315	A1	31-12-1998
			JP	1299827	Α	04-12-1989
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			US	5063254	Α	05-11-1991
US 2002031488	A1	14-03-2002	JP	2002097366	A	02-04-2002

PATENT COOPERATION TREATY

PCT

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY (Chapter I of the Patent Cooperation Treaty)

(PCT Rule 44bis)

Applicant's or agent's file reference DC1009 PCT 1	FOR FURTHER ACTION	See item 4 below			
International application No. PCT/US2005/002451	International filing date (day/month/year) 20 January 2005 (20.01.2005)	Priority date (day/month/year) 02 February 2004 (02.02.2004)			
International Patent Classification (8th edition unless older edition indicated) See relevant information in Form PCT/ISA/237					
Applicant DOW CORNING CORPORATION					

1.	This international preliminary report on patentability (Chapter I) is issued by the International Bureau on behalf of the International Searching Authority under Rule 44 bis.1(a).						
2.	This REPORT consists of a total	of 6 sheets, including this cov	ver sheet.				
٠	In the attached sheets, any refere to the international preliminary re	nce to the written opinion of t eport on patentability (Chapte	he International Searching Authority should be read as a reference r I) instead.				
3.	This report contains indications r	elating to the following items					
	Box No. I	Basis of the report					
	Box No. II	Priority					
	Box No. III	Non-establishment of opini applicability	on with regard to novelty, inventive step and industrial				
	Box No. IV	Lack of unity of invention					
	Box No. V	Reasoned statement under applicability; citations and	Article 35(2) with regard to novelty, inventive step or industrial explanations supporting such statement				
	Box No. VI	Certain documents cited					
	Box No. VII	Certain defects in the intern	national application				
	Box No. VIII	Certain observations on the	international application				
4.	4. The International Bureau will communicate this report to designated Offices in accordance with Rules 44bis.3(c) and 93bis.1 but not, except where the applicant makes an express request under Article 23(2), before the expiration of 30 months from the priority date (Rule 44bis.2).						
	Date of issuance of this report 07 August 2006 (07.08.2006)						
	The International Burea 34, chemin des Colo 1211 Geneva 20, Sw	ombettes	Authorized officer Philippe Becamel				
Facsir	mile No. +41 22 338 82 70		e-mail: pt12@wipo.int				

Form PCT/IB/373 (January 2004)

PATENT COOPERATION TREATY

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				(P	CT Rule 43bis.1)
				Date of mailing	
				(day/month/year) see	form PCT/ISA/210 (second sheet)
	- 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1			FOR FURTHER A	CTION
	icant's or agent's file reform PCT/ISA/22			See paragraph 2 below	N
	national application N		International filing date (day/month/year)	Priority date (day/month/year)
	T/US2005/002451	o .	20.01.2005		02.02.2004
		Ification (IPC) or	both national classification	and IPC	
CO	8G77/44, C08G77	04, A61K7/11	, C08G77/10		
App	licant				
DC	W CORNING CO	RPORATION			
1.	This opinion co	ntains indicati	ons relating to the fol	lowing items:	
	⊠ Box No. I	Basis of the or	oinion		•
	☐ Box No. II	Driority			
	☐ Box No. III	Non-establish	ment of opinion with reg	ard to novelty, inventi	ve step and industrial applicability
ŀ	☐ Box No. IV	Lack of unity of	of invention		the inventive stop or industrial
	☑ Box No. V	Reasoned sta applicability; o	tement under Rule 43 <i>b.</i> itations and explanation	<i>is</i> .1(a)(i) with regard to as supporting such sta	novelty, inventive step or Industrial tement
	☐ Box No. VI	Certain docum			
1	□ Box No. VII		ts in the international ap		
	☐ Box No. VIII	Certain obser	vations on the internation	onal application	
2.	FURTHER ACT	ION			
	written opinion of the applicant che International Bu will not be so co	of the Internation ooses an Author reau under Rula Insidered.	ority other than this one e 66.1 <i>bis</i> (b) that written	to be the IPEA and the opinions of this Intern	Il usually be considered to be a However, this does not apply where s chosen IPEA has notifed the ational Searching Authority
	If this opinion is, submit to the IP months from the whichever expir	EA a written reported the second control of	pove, considered to be a ply together, where app g of Form PCT/ISA/220	a written opinion of the ropriate, with amendm or before the expiration	PPEA, the applicant is invited to lents, before the expiration of three n of 22 months from the priority date,
	For further optic	ons, see Form F	PCT/ISA/220.		
3.	For further deta	ils, see notes to	Form PCT/ISA/220.		
	-				
- 1					

Name and mailing address of the ISA:

<u>)</u>

European Patent Office D-80298 Munich Tel. +49 89 2399 - 0 Tx: 523656 epmu d Fax: +49 89 2399 - 4465 Authorized Officer

Buestrich, R

Telephone No. +49 89 2399-7473



WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No. PCT/US2005/002451

	Box I	
1.	the la	regard to the language , this opinion has been established on the basis of the international application in nguage in which it was filed, unless otherwise indicated under this item.
	□ 1 l: (This opinion has been established on the basis of a translation from the original language into the following anguage , which is the language of a translation furnished for the purposes of international search under Rules 12.3 and 23.1(b)).
2.	With neces	regard to any nucleotide and/or amino acid sequence disclosed in the international application and ssary to the claimed invention, this opinion has been established on the basis of:
	a. typ	pe of material:
		a sequence listing
		table(s) related to the sequence listing
	b. fo	rmat of material:
		l in written format
		in computer readable form
	c. tin	ne of filing/furnishing:
		filed together with the international application in computer readable form.
		furnished subsequently to this Authority for the purposes of search.
3	3. 🗆	In addition, in the case that more than one version or copy of a sequence listing and/or table relating therein has been filed or furnished, the required statements that the information in the subsequent or additional copies is identical to that in the application as filed or does not go beyond the application as filed, as appropriate, were furnished.
	4. Add	litional comments:

Box No. V Reasoned statement under Rule 43bis.1(a)(i) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. Statement

Novelty (N)

Yes: Claims

1-7

No:

Claims

Inventive step (IS)

Yes: Claims

1-7

Claims No:

Industrial applicability (IA)

Yes: Claims

1-7

No: Claims

2. Citations and explanations

see separate sheet

Re item V:

The following documents are referred to in this communication: 1

D1: EP-A-0 336 633 (LOCTITE CORPORATION) 11 October 1989 (1989-10-11)

D2: US 2002/031488 A1 (KANJI MOHAMED ET AL) 14 March 2002 (2002-03-14)

The present application meets the criteria of Article 33(1) PCT, because the subject-2 matter of independent claim 1 is new in the sense of Article 33(2) PCT.

Document D1 discloses MTQ siloxane resin comprising the units (D1, example 1):

- $(R^{1}_{3}SiO_{1/2})_{a}$
- (iii) $(R^3SiO_{3/2})_c$
- (iv) $(SiO_{4/2})_d$

wherein

a=0.4, c=0.12 and d=0.48.

The subject-matter of claim 1 of the present application differs from D1 in that greater than 40 mol% of the R3 groups are propyl groups.

Therefore the subject-matter of claim 1 is novel over D1.

Document D2 discloses cosmetic compositions comprising siloxane resin comprising MQ-resins and T-resins (D2, paragraph 0016 and example 1).

The subject-matter of claim 1 of the present application differs from D2 in that a homogeneous MTQ-resin is required.

Therefore the subject-matter of claim 1 is novel over D2.

The present application meets the criteria of Article 33(1) PCT, because the subject-3 matter of independent claim 1 is inventive in the sense of Article 33(3) PCT.

D2 discloses similar compositions for cosmetic applications and is therefore regarded as closest prior art.

The objective technical problem of the present application is to provide a siloxane resins which show comparable extended wear and transfer resistance and improved gloss and which are non-tacky. It is shown in the examples (cf. table 6) that the MTQ-resins claimed, combine the positive properties of MQ and T-resins.

There is no indication in the prior art, including D1 and D2, to synthesize MTQ-resins for cosmetic compositions which show comparable extended wear and transfer resistance and improved gloss and which are non-tacky.

The subject-matter of claim 1 is therefore inventive (Article 33(3) PCT).

- The subject-matter of present independent claims 3, 4, 5 refer to a method of making said MTQ-resin, a siloxane resin prepared by the method and a personal care product comprising said resin. Since the resin as such is novel and inventive these claims are as well novel and inventive.
- 5 Claims 2, 6 and 7 are dependent claims and as such also meet the requirements of the PCT with respect to novelty and inventive step.

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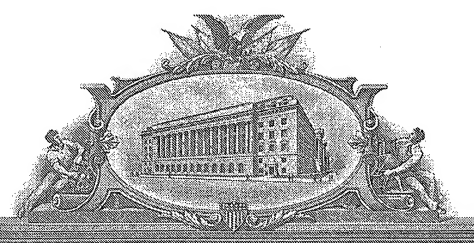
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March 02, 2005

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Respectfully submitted

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PROVISIONAL APPLICATION FOR PATENT COVER SHEET This is a request for rilling a PROVISIONAL APPLICATION FOR PATENT und r 37 CFR 1.53(c).

INVENTOR(S) Residence Family Name or Surname (City and either State or Foreign Country) Given Name (first and middle [if any]) Turner, Michigan Cook Julie Lyn Midland, Michigan Hinterman Daniel M. Stark-Kasley Midland, Michigan L ri Ann Midland, Michigan Gary Michael Wieber separately numbered sheets attached hereto Additional inventors are being named on the TITLE OF THE INVENTION (280 characters max) MQ - T Propyl Siloxane Resins **CORRESPONDENCE ADDRESS** Direct all correspondence to: Place Customer Number 00137 **Customer Number** Bar Code Label here OR Type Customer Number here Firm or Individual Name <u>Address</u> Address State ZIP City Country **ENCLOSED APPLICATION PARTS (check all that apply)** Specification Number of Pages CD(s), Number Drawing(s) Number of Sheets Claims - 3 pages Other (specify) Abstract - 1 page Application Data Sheet. See 37 CFR 1.76 METHOD OF PAYMENT OF FILING FEES FOR THIS PROVISIONAL APPLICATION FOR PATENT (check one) **FILING FEE** AMOUNT (\$) A check or money order is enclosed to cover the filing fees The Director is hereby authorized to charge filing 04-1520 \$160.00 fees or credit any overpayment to Deposit Account Number Payment by credit card. Form PTO-2038 is attached. The invention was made by an agency of the United States Government or under a contract with an agency of the United States Government. X No. Yes, the name of the U.S. Government agency and the Government contract number are:

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This collection of information is required by 37 CFR 1.51. The information is used by the public to file (and by the PTO to process) a provisional application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 8 hours to complete, including gathering, preparing, and submitting the complete provisional application to the PTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Mail Stop Provisional Application, Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.



UNITED STATES PATENT APPLICATION FOR

MQ - T Propyl Siloxane Resins

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Having the inventor(s):

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Julie Lyn Cook
Daniel M. Hinterman
Lori Ann Stark-Kasley
Gary Michael Wieber

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FIELD OF THE INVENTION

The present invention provides siloxane resins, herein known as MQ-T propyl resins, comprising (R¹₃SiO_{1/2})_a, (R²₂SiO_{2/2})_b, (R³SiO_{3/2})_c, and (SiO_{4/2})_d units, where at least 40 mole % of the R³ groups are propyl. The present invention further relates to a method of preparing such siloxane resins by reacting a MQ siloxane resin with a T propyl siloxane resin. The siloxane resins of the present invention are useful in a variety of personal, household, and medical care applications, and in particular, as a resin additive in pigmented cosmetic formulations.

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BACKGROUND OF THE INVENTION

Siloxane resins of the general formula R_nSiO_{(4-n)/2}, where R is an alkyl group and n is generally less than 1.8, are an important family of silicone polymers because of their utility in many commercial applications such as adhesive compositions and coatings applications. One particular subclass of siloxane resins, known as MQ resins (since they comprise primarily of "M" units of the general formula R₃SiO_{1/2} and "Q" units of the general formula SiO₂), have found utility in cosmetic formulations. In particular MQ resins are commonly used in "extended wear" or "transfer resistant" cosmetic formulations. In these formulations, MQ are resins enhance the substantivity of the pigments or other formulation actives to skin after application creating a longer lasting, and hence extended wear product.

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Representative examples of transfer resistant cosmetic compositions using MQ resins are found in US 6,071,503, US 6,074,654, US 6,139,823, US 6,340,466, WO 97/17058, and WO 97/17059 which disclose compositions comprising the combination of organosiloxane resins and fluid diorganosiloxane resins with a volatile carrier.

US 5,330,747 teaches cosmetics with enhanced durability using a film forming agent from a pressure sensitive adhesive composition comprising a trimethylsilyl endblocked resinous copolymer, a silanol endblocked polydiorganosiloxane fluid, and a phenyl containing polysiloxane fluid.

US 5, 075,103 and US 5,733,537 teach a hair treating method for imparting curl retention to hair in which at least one film forming ingredient is applied to the hair. The improvement utilizes as the film forming ingredient an organosilicon compound which is a nonpolar silsequioxane.

US 5,800,816 discloses cosmetic compositions having improved transfer resistance, comprising: a) from about 0.1-60% by weight of trimethylated silica, b) from about 0.1-60% by weight of a volatile solvent having a viscosity of 0.5 to 100 centipoise (mPa·s) at 25°C, c) 0.1-60% by weight of a nonvolatile oil having a viscosity of 200 to 1,000,000 centipoise (mPa·s) at 25°C, d) 0.1-80% of a cosmetically acceptable carrier.

US 5,837,223 and US 6,036,947 teach transfer resistant high luster cosmetic stick compositions comprising, by weight of the total composition: a) 10-70% of a volatile solvent

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having a viscosity of 0.5 to 20 centipoise (mPa·s) at 25.degree. C., b) 0.5-40% of a guerbet ester, and c) 0.1-20% of a siloxysilicate polymer.

GB 2,319,527 discloses fragrance releasing non-volatile polysiloxanes based on a high molecular weight polydiorganosiloxane compounds where at least one or more of the organic substituents of the polymer is a radical derived from a fragrant alcohol.

Japanese examined patent publication 1994-72085 teaches makeup cosmetic: compositions having improved water resistance and durability containing an organic silicone resin, a volatile silicone oil, and a make up powder.

While the use of MQ resins in cosmetics have led to formulations having extended wear or transfer resistance, a need exists to alter the properties of the siloxane resins used in such formulations. In particular, films of MQ resins used in these formulations can have a matte finish and feel tacky. Thus, there is a need for improved siloxane resins that offer at least comparable extended wear and transfer resistance properties of the MQ resins presently used in cosmetic formulations, but having improved gloss (i.e. non-matte) that are non-tacky. Furthermore, there is a need for resins in hair care formulations that improve the curl retention properties of hair following treatment.

The present inventors have discovered improved siloxane resins by incorporating propyl containing siloxane (R³SiO_{3/2})_c units (T units) with M and Q units. The resulting siloxane resins, herein referred to as MQ – T propyl siloxane resins, have improved physical properties. In particular, cosmetic formulations containing the present MQ-T propyl siloxane resins have improved gloss vs MQ resins and are less tacky than T propyl resins, while maintaining their long lasting or wear characteristics.

SUMMARY OF THE INVENTION

This invention relates to a MQ-T propyl siloxane resin comprising the units:

(i) $(R^1_3SiO_{1/2})_a$

(ii) $(R^2_2SiO_{2/2})_b$

(iii) (R3SiO3/2)c, and

(iv) (SiO_{4/2})_d

wherein

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R¹, R², and R³ are independently an alkyl group having from 1 to 8 carbon atoms, an aryl group, a carbinol group, or an amino group,

a has a value 0.05 to 0.5,

b has a value of zero to 0.3,

c has a value greater than zero,

d has a value of 0.05 to 0.6,

the value of a + b + c + d = 1,

with the proviso that greater than 40 mole% of the R³ groups in the siloxane resin are propyl.

The invention further relates to a method of making a siloxane resin composition and the products obtained therefrom. The method comprises reacting:

A) a MQ resin comprising at least 80 mole % (R¹₃SiO_{1/2})_a and (SiO_{4/2})_d units where R¹ is an alkyl group having from 1 to 8 carbon atoms, an aryl group,

a carbinol group, or an amino group,

a and d has a value greater than zero,

the ratio of a/d is 0.5 to 1.5;

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B) a T propyl resin comprising at least 80 mole % R3SiO units,

where R3 is an alkyl group having from 1 to 8 carbon atoms, an aryl group,

a carbinol group, or an amino group,

c has a value greater than zero,

and with the provisio that at least 40 mole % of the R³ groups are propyl, wherein the weight ratio of A/B is from 95:5 to 15:85.

The MQ-T propyl siloxane resins are useful in a variety of personal, household, or medical care compositions. In particular, the MQ-T propyl siloxane resins provide glossy, non-tacky films that can be used to enhance the substantivity of color cosmetic formulations. The MQ-T propyl siloxane resins can also be used as additives in hair care formulations to enhance curl retention properties. The MQ-T propyl siloxane resins can also enhance the feel of treated hair, providing a softer feel compared to other siloxane resin additives. Thus, the present invention provides personal, household, or medical care compositions comprising the MQ-T propyl siloxane resins described herein.

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DETAILED DESCRIPTION OF THE INVENTION

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The MQ-T propyl siloxane resin of the present invention comprises (i) $(R^1 _3 SiO_{1/2})_a$, (ii) $(R^2 _2 SiO_{2/2})_b$, (iii) $(R^3 SiO_{3/2})_c$, and (iv) $(SiO_{4/2})_d$ units which are commonly known in the art, and also used herein, as M, D, T, and Q units respectively. The amount of each unit present in the MQ-T propyl siloxane resin can be expressed as a mole fraction (i.e. a, b, c, or d) of the total number of moles of all M, D, T, and Q units present in the MQ-T propyl siloxane resin. The value of a (mole fraction of M units) is 0.05 - 0.5, or alternatively 0.15 to 0.4. The value of b (mole fraction of D units) is 0 - 0.3, alternatively 0 to 0.1, or alternatively 0 to 0.05. Thus, the MQ-T propyl siloxane resins can be free of D units, or alternatively can contain up to and including 0.3 mole fraction of D units. The value of c (mole fraction of T units) is greater than 0, alternatively 0.05 to 0.65, or alternatively 0.4 to 0.65. The value of d (mole fraction of Q units) is 0.05 to 0.6, alternatively 0.2 to 0.6, or alternatively 0.2 to 0.55:

The MQ-T propyl siloxane resins of the present invention are characterized by having at least 40 mole %, alternatively 50 mole %, or alternatively 90 mole % of the R³ alkyl groups on the T unit be propyl.

The R¹, R², and R³ in the units of the MQ-T propyl siloxane resin are independently an alkyl group having from 1 to 8 carbon atoms, an aryl group, a carbinol group, or an amino group. The alkyl groups are illustrated by methyl, ethyl, propyl, butyl, pentyl, hexyl, and octyl. The aryl groups are illustrated by phenyl, naphthyl, benzyl, tolyl, xylyl, xenyl, methylphenyl, 2-phenylethyl, 2-phenyl-2-methylethyl, chlorophenyl, bromophenyl and fluorophenyl with the aryl group typically being phenyl.

For the purposes of this invention a "carbinol group" is defined as any group containing at least one carbon-bonded hydroxyl (COH) radical. Thus the carbinol groups may contain more than one COH radical such as for example

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The carbinol group if free of aryl groups has at least 3 carbon atoms, or an aryl-containing carbinol group having at least 6 carbon atoms, The carbinol group free of aryl groups having at least 3 carbon atoms is illustrated by groups having the formula R⁴OH wherein R⁴ is a divalent hydrocarbon radical having at least 3 carbon atoms or divalent hydrocarbonoxy radical having at least 3 carbon atoms. The group R⁴ is illustrated by alkylene radicals such as -(CH₂)_x- where x has a value of 3 to 10,
-CH₂CH(CH₃)-, -CH₂CH(CH₃)CH₂-, -CH₂CH₂CH(CH₂CH₃)CH₂CH₂CH₂-, and
-OCH(CH₃)(CH₂)_x- wherein x has a value of 1 to 10.

The aryl-containing carbinol group having at least 6 carbon atoms is illustrated.

10 by groups having the formula R⁵OH wherein R⁵ is an arylene radical such as

-(CH₂)_xC₆H₄- wherein x has a value of 0 to 10, -CH₂CH(CH₃)(CH₂)_xC₆H₄- wherein x has a value of 0 to 10, -(CH₂)_xC₆H₄(CH₂)_x- wherein x has a value of 1 to 10. The aryl
containing carbinol groups typically have from 6 to 14 atoms.

The amino group is illustrated by groups having the formula $-R^6NH_2$ or $-R^6NHR^7NH_2$ wherein R^6 is a divalent hydrocarbon radical having at least 2 carbon atoms and R^7 is a divalent hydrocarbon radical having at least 2 carbon atoms. The group R^6 is typically an alkylene radical having from 2 to 20 carbon atoms. R^6 is illustrated by ethylene, propylene, $-CH_2CHCH_3$ -, butylene, $-CH_2CH(CH_3)CH_2$ -, pentamethylene, hexamethylene, 3-ethyl-hexamethylene, octamethylene, and decamethylene.

R⁷ is typically an alkylene radical having from 2 to 20 carbon atoms. R⁷ is illustrated by ethylene, propylene, -CH₂CHCH₃-, butylene, -CH₂CH(CH₃)CH₂-, pentamethylene, hexamethylene, 3-ethyl-hexamethylene, octamethylene, and decamethylene.

Typical amino groups are -CH₂CH₂CH₂NH₂ and -CH₂(CH₃)CHCH₂(H)NCH₃, -CH₂CH₂NHCH₂CH₂NHCH₂, -CH₂CH₂NHCH₃, -CH₂CH₂CH₂CH₂NHCH₃, -(CH₂CH₂NHCH₃), and -CH₂CH₂NHCH₂NHCH₂NHCH₃.

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Typically, R^1 is a methyl group, R^2 is a methyl or phenyl group, and R^3 is a propyl group.

Any individual D, T or Q siloxane units of the MQ-T propyl siloxane resins can also contain a hydroxy group and/or alkoxy group. Such siloxane units containing hydroxy and/or alkoxy groups are commonly found in siloxane resins having the general formula R_nSiO_{(4-n)/2}. The hydroxy groups in these siloxane resins typically result from the reaction of the hydrolyzable group on the siloxane unit with water; the alkoxy groups result from incomplete hydrolysis when alkoxysilane precursors are used or from exchange of alcohol with hydrolysable groups. Typically, the weight percent of the total hydroxy groups present in the MQ-T propyl siloxane resin is 3 %, alternatively, 2 %, or alternatively, 1.5%. Typically, the weight percent of the total alkoxy groups present in the MQ-T propyl siloxane resin is up to 20 %, alternatively up to 10 %.

The molecular weights of the MQ-T propyl siloxane resins are not restricted, but typically the number average molecular weight (M_N) ranges from 3,000 to 10,000, or alternatively from 5,000 to 8,000.

The MQ-T propyl siloxane resins of the present invention can be prepared by any of the methods known in the art for preparing siloxane resins having the general formula $R_n SiO_{(4-n)/2}$ where R is an alkyl group and n is generally less than 1.8. Alternatively, the MQ-T propyl resins can be prepared by the methods described infra.

The MQ-T propyl resins of this invention are illustrated by:

MQ-T propyl resins comprising the units;

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((CH_3)_3SiO_{1/2})_a, (R^3SiO_{3/2})_c, where R^3=CH_3CH_2CH_2-, and (SiO_{4/2})_d
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25 MQ-T propyl resins comprising the units;

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\begin{split} &((CH_3)_3SiO_{1/2})_a\;,\\ &((CH_3)_2SiO_{2/2})_b\;,\\ &(R^3SiO_{3/2})_c\;, \text{ where } R^3=CH_3CH_2CH_2\text{--, and}\\ &(SiO_{4/2})_d \end{split}
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30 MQ-T propyl resins comprising the units;

 $((CH_3)_3SiO_{1/2})_a$,

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((CH_3)_2SiO_{2/2})_b, ((CH_3)(C_6H_5)SiO_{2/2})_{b'},
              (R^3SiO_{3/2})_c, where R^3 = CH_3CH_2CH_2-, and
               (SiO_{4/2})_{d}
      MQ-T propyl resins comprising the units;
               ((CH_3)_3SiO_{1/2})_a,
               ((CH_3)_2SiO_{2/2})_b,
              (R^3SiO_{3/2})_c, where R^3 = CH_3CH_2CH_2-, and (C_6H_5SiO_{3/2})_c
               (SiO<sub>4/2</sub>)<sub>d</sub>
      MQ-T propyl resins comprising the units;
               ((CH_3)_3SiO_{1/2})_a,
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               ((CH_3)_2SiO_{2/2})_b, ((CH_3)(C_6H_5)SiO_{2/2})_{b'},
               (R^3SiO_{3/2})_c, where R^3 = CH_3CH_2CH_2-, (C_6H_5SiO_{3/2})_c, and
               (SiO_{4/2})_d
               wherein a has a total value in the resin of 0.05 to 0.5, the sum of b + b' has a total
      value in the resin of zero to 0.3, c has a total value in the resin of 0.05 to 0.65, and d has a
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      total value in the resin of 0.05 to 0.6.
               The present invention also provides a method of making a siloxane resin composition
      and the products obtained therefrom. The method comprises reacting:
               A) a MQ resin comprising at least 80 mole % (R^{1}_{3}SiO_{1/2})_{a} and (SiO_{4/2})_{d} units
                       where R<sup>1</sup> is an alkyl group having from 1 to 8 carbon atoms, an aryl group,
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                                a carbinol group, or an amino group,
                                a and d has a value greater than zero,
                                the ratio of a/d is 0.5 to 1.5;
      and
               B) a T propyl resin comprising at least 80 mole % R3SiO units,
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                       where R<sup>3</sup> is an alkyl group having from 1 to 8 carbon atoms, an aryl group,
                                a carbinol group, or an amino group,
                                c has a value greater than zero,
                       and with the provisio that at least 40 mole % of the R<sup>3</sup> groups are propyl,
       wherein the weight ratio of A/B is from 95:5 to 15:85.
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Component A) is a MQ resin comprising at least 80 mole % (R¹₃SiO_{1/2})_a and (SiO_{4/2})_d units where R¹ is the same as defined above, i.e. an alkyl group having from 1 to 8 carbon atoms an aryl group, a carbinol group, or an amino group, a and d has a value greater than zero, and the ratio of a/d is 0.5 to 1.5. MQ resins suitable for use as component (A), and methods for their preparation, are known in the art. For example, U.S. Patent No. 2,814,601 to Currie et al., November 26, 1957, which is hereby incorporated by reference, discloses that MQ resins can be prepared by converting a water-soluble silicate into a silicic acid monomer or silicic acid oligomer using an acid. When adequate polymerization has been achieved, the resin is end-capped with trimethylchlorosilane to yield the MQ resin. Another method for preparing MQ resins is disclosed in U.S. Patent No. 2,857,356 to Goodwin, October 21, 1958, which is hereby incorporated by reference. Goodwin discloses a method for the preparation of an MQ resin by the cohydrolysis of a mixture of an alkyl silicate and a hydrolyzable trialkylsilane organopolysiloxane with water.

The MQ resins suitable as component A) in the present invention may contain D and T units, providing that at least 80 mole %, alternatively 90 mole % of the total siloxane units are M and Q units. The MQ resins may also contain hydroxy groups. Typically, the MQ resins have a total weight % hydroxy content of 2-10 weight %, alternatively 2-5 weight.%. The MQ resins can also be further "capped" wherein residual hydroxy groups are reacted further with M groups.

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Component B) is a T propyl resin comprising at least 80 mole % of R³SiO_{3/2} units, where R³ is the same as defined above, i.e. an alkyl group having from 1 to 8 carbon atoms an aryl group, a carbinol group, or an amino group, and with the proviso that at least 40 mole % of the R³ groups are propyl. Typically, the T propyl resin is a silsesquioxane resin.

Silsesquioxane resins are well known in the art and are typically prepared by hydrolyzing an organosilane having three hydrolyzable groups, such as a halogen or alkoxy group, present in the molecule. Thus, component (B) can be obtained by hydrolyzing propyltrimethoxysilane, propyltriethoxysilane, propyltripropoxysilane, or by co-hydrolyzing the aforementioned propylalkoxysilanes with various alkoxysilanes. Examples of these alkoxysilanes include methyltrimethoxysilane, methyltriethoxysilane, methyltriisopropoxysilane, dimethyldimethoxysilane, and phenyltrimethoxysilane. Propyltrichlorosilane can also be hydrolyzed alone, or in the presence of alcohol. In this case, co-hydrolyzation can be carried

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out by adding methyltrichlorosilane, dimethyldichlorosilane, phenyltrichlorosilane, or similar chlorosilanes and methyltrimethoxysilane, methyltriethoxysilane, methyltriisopropoxysilane, or similar methylalkoxysilane. Alcohols suitable for these purposes include methanol, ethanol, n-propyl alcohol, isopropyl alcohol, butanol, methoxy ethanol, ethoxy ethanol, or similar alcohols. Examples of hydrocarbon-type solvents which can also be concurrently used include toluene, xylene, or similar aromatic hydrocarbons; hexane, heptane, isooctane, or similar linear or partially branched saturated hydrocarbons; and cyclohexane, or similar aliphatic hydrocarbons.

The T-propyl resins suitable as component B) in the present invention may contain M, D, and Q units, providing that at least 80 mole %, alternatively 90 mole % of the total siloxane units are T units. The T-propyl resins may also contain hydroxy groups. Typically, the T-propyl resins have a total weight % hydroxy content of 3-8 weight %.

A polyorganosiloxane can optionally be added to the method of the present invention as component C). Polyorganosiloxane useful as component C) in the present invention comprise R²₂SiO_{2/2}, (i.e. D units) or R³SiO_{3/2} (T units). The polyorganosiloxane can be added to introduce various D and T units into MQ-T propyl resins to alter the properties of the resulting resins. The structure or formula of the polyorganosiloxane is not restrictive, e providing the polyorganosiloxane contains some measurable quantity of R²₂SiO_{2/2}, units (i.e. D units) or R³SiO_{3/2} (T units), and the total amount of polyorganosiloxane added to the reaction of A) and B) does not provide more than 50 mole % D or T units into the reaction mixture. The polyorganosiloxane can contain any combination of M, D, T and Q units, providing at least some D or T units are present. Thus, the polyorganosiloxane can be selected from any of the fluid, gum, or resinous silicones known in the art containing D or T units, or combinations thereof. The D units typically contain methyl or phenyl as the R² substituents, which can be designated as D^{Me} and D^{Ph} respectively, or any combinations thereof. The T units typically contain methyl or phenyl as the R³ substituents, which can be designated as T^{Me} and T^{Ph} respectively, or any combinations thereof. The polyorganosiloxane can be a linear polydiorganosiloxane fluid having a viscosity of 10 -1000 cS (mm²/s). Typically the polydiorganosiloxane fluid is polydimethylsiloxane, or alternatively a polymethylphenylsiloxane. The polyorganosiloxane can also be an

organosilsequioxane resin. The organosilsequioxane resin typically is a methylsilsequioxane resin or a phenylsilsequixone resin.

Components A), B), and optionally C) can be reacted by any method in the art known to effect reaction of M, D, T, and Q siloxane units. Typically however, components A), B), and optionally C) are reacted by a condensation reaction in the presence of a catalyst. Typically the MQ resin is contained in an aromatic hydrocarbon or siloxane solvent. Suitable condensation reaction catalysts are base catalysts including metal hydroxides such as potassium hydroxide and sodium hydroxide; metal salts such as silanolates, carboxylates, and carbonates; ammonia; amines; and titanates such as tetrabutyl titanates; and combinations thereof. Typically the reaction of components A), B), and optionally C) is effected by heating the reaction mixture to temperatures ranging from 50 to 140°C, alternatively 100 to: 140°C. The reaction can be conducted in a batch, semi-continuous, or continuous process.

The weight ratio of component A) to component B) (i.e. A/B) in the reaction can vary from 95:5 to 15:85, alternatively 95:5 to 20:80, or alternatively 90:10 to 20:80. The amount of component C) can vary, but the amount of component C) added should introduce less than 30 mole% of additional D or T units, based on the total moles of siloxane units in the reaction mixture.

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The MQ-T propyl resins of the present invention are useful in a variety of personal, household, or medical care applications. In particular, the MQ-T propyl siloxane resins provide glossy, non-tacky films that can be used to enhance the substantivity of color cosmetic formulations. The MQ-T propyl siloxane resins can also be used as additives in hair care formulations to enhance curl retention properties. Thus, the present invention provides personal, household, or medical care compositions comprising the MQ-T propyl siloxane resins described herein.

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EXAMPLES

The following examples are presented to further illustrate the compositions and methods of this invention, but are not to be construed as limiting the invention. All parts and percentages in the examples are on a weight basis and all measurements were obtained at about 23°C, unless indicated to the contrary.

Materials

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MQ Resin = a MQ resin having the formula $M_{0.43}Q_{0.57}$ and M_n = 3230 dissolved in xylene at 70.8 wt % solids. The MQ resin was prepared according to techniques taught by Daudt in US 2,676,182.

T propyl resin = propyl silsesquioxane resin at 74.8 wt % in toluene. The propyl silsesquioxane resin was prepared from the hydrolysis of propyl trichlorosilane.

Phenyl silsesquioxane resin = a phenyl silsesquioxane solid flake resin at 100 wt % solids prepared from the hydrolysis of phenyl trichlorosilane.

Example 1 Preparation of MQT^{Pr} Resins

A 3-neck reaction flask equipped with an agitator, temperature probe, and a Dean Stark trap fitted with a condenser on top, was loaded with MQ Resin, T propyl resin, xylene and 1M KOH in water at the ratios as shown in Table 1. The Dean Stark trap was pre-loaded with xylene to insure a 50% solids level in the reaction flask was maintained. The mixture in the flask was maintained at reflux temperature (100-140°C) for at least 3 hours. Any water that formed in the reaction mixture was continuously removed as needed and trapped as an azeotrope in the Dean Stark trap. After 3 hours of reflux, water was removed from the trap, and heating continued for an additional 30 min. After allowing the mixture to cool, excess acetic acid was added to neutralize the KOH in the mixture. The mixture was then filtered to remove the salts formed by passing it through a pressure filter. A solvent exchange was completed by heating the mixture in a rotary evaporator under vacuum. After the majority of

xylene was removed, decamethylcyclopentasiloxane was added while continuing to remove any residual aromatic solvent. The structures of the resulting siloxane resins were characterized by ²⁹Si NMR spectroscopy and GPC and the results summarized in Table 4 below.

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Table 1

Example #	Wt Ratio MQ/T ^{PT} resins added	Wt% MQ Resin	Wt% T Propyl Resin	Wt% Xylene	Wt% IM KOH	Wt% Acetic Acid
l-a	(85:15)	59.4	10.5	29.1	0.9	0.2
1- b	(50:50)	34.9	34.8	29.1	0.9	0:2
1- c	(30:70)	20.9	48.8	29.2	0.9	0.2
1-d	(95:5)	67.1	3.5	28.3	0.9	0.2
1- e	(100:0)	69.3	0	28.8	0.9	0.2

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Example 2

Preparation of $MQT^{Pr}T^{Ph}$ resins

A series of MQT^{Pr}T^{Ph} resins were prepared following the experimental procedure described in Example 1 using the formulations shown in Table 2. In this series, a phenyl silsesquioxane resin was added to incorporate additional phenyl containing T units into the siloxane resin. The structures of the resulting siloxane resins were characterized by ²⁹Si NMR spectroscopy and GPC, as summarized in Table 4 below.

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Table 2

Example #	Wt Ratio MQ/T ^{Pr} /T ^{Ph} resins	Wt% MQ Resin	Wt% T Propyl Resin	Wt% T Phenyl Resin	Wt% of Xylene	Wt of 1M KOH solution in water	Wt of Glacial Acetic Acid
2-a	(85:7.5:7.5)	59.4	5.2	3.7	30.6	0.9	0.2
2-b	(50:25:25)	34.7	17.4	12.4	34.4	0.9	0.2
2-c	(15:42.5:42.5)	10.5	29.6	21.0	37.8	0.9	0.2

25 Example 3

Preparation of MQT^{Pr}D resins

Two MQT^{Pr}D resins were prepared following the experimental procedure described in Example 1 using the formulations shown in Table 3. In this series, a 100 cst

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polydimethylsiloxane fluid was added to incorporate D units (Me₂SiO) into the siloxane resin. The structures of the resulting siloxane resins were characterized by ²⁹Si NMR spectroscopy and GPC, as summarized in Table 4 below.

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Table 3

Example #	Wt Ratio MQ/T ^{Pt} /D added	Wt% MQ Resin	Wt% T Propyl Resin	Wt% D	Wt% of Xylene	Wt of 1M KOH solution in water	Wt of Glacial Acetic Acid
3-a	(38:57:5)	26.9	39.7	2.5	29.9	0.9	0.2
3-b	(28.5:66.5:5)	14.1	32.9	2.5	30.0	0.9	0.2

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Table 4

Example #	Resin structure according to NMR Characterization	wt% OH	Mn	Mw	Mw/Mn
MQ resin	M ^{0.43} Q ^{0.57}		3230	1516	4.7
T propyl resin	T ^{Pr} _{1,0}	7.0	3470	11400	3.3
l-a	M _{0.374} Q _{0.529} :T ^{PT} _{0.097}	1.4	5880	271000	46.1
1-b	$M_{0.248}Q_{0.341}:T^{Pr}_{0.412}$	2.1	6640	3860000	581.3
1-c	$M_{0.162}Q_{0.217}$: $T^{Pr}_{0.621}$	1.5	7600	25300000	3329
1-d	M _{0.419} Q _{0.5485} ; T ^{PT} _{0.03}	1.5			
1-e	M Q	1.7	5200	28900	5.6
2-a	$M_{0.396}Q_{0.523}:T^{PT}_{0.0491}:T^{Ph}_{0.0339}$	1.6	5200	26900	5.2
2 - b	M _{0.272} Q _{0.347} :T ^{PT} _{0.220} :T ^{Ph} _{0.161}	0.8	5867	1640000	279
2-с	M 0.09 Q0 118: TPT 0.457: TPD 0.334	0.8			
3-a	MQ:T ^{PT} :D ^{Me2}		5455	303000	55.5
3-b	$M_{0.155}Q_{0.201}$: $T^{Pr}_{0.591}$: $D^{Me2}_{0.053}$	1.8	5140	295000	57.4

Films resulting from the siloxane resins of Examples 1-3 were evaluated for gloss and tack after coating from 35% solids in volatile solvent onto a Leneta chart, evaluated in a foundation (color cosmetic) for durability, and in a hair fixative for curl retention and conditioning. The results are summarized in Table 6. The siloxane resins were applied to the hair tresses as 6 wt. % solutions in decamethylcyclopentasiloxane or isododecane.

Descriptions of the specific test methods and composition of the cosmetic formulation used in these evaluations follows Table 6.

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Table 6

Example #	60° Gloss	Tack	Foundation Durability - ΔE (Change in color)	Curl Retention – Hair – % curl maintained after 5 hours at 95% RH	Hair Conditioning - Wet Comb % Reduction in combing force	Hair Conditioning – Dry Comb % Reduction in combing force
No Resin – neg control	55 blank chart		9.2	30.5 -36.8	-95	-50
MQ resin	34.8	Not tacky	3.8	53-58	-55.7	-1650
T propyl resin	78.4	Very tacky	11.0	44		
l-a	72.6	Not Tacky	5.1	45.5		•
1-b	74.5	slight, less w/time	4.7	50.6	-65	-1648
1-c	75.2	slight, less w/time	5.4	51.2	2.2	-1508
1-d	68.2	Not tacky				.,
l-e	68.7	Not tacky				
2-a	62.3	Not Tacky	3.1	41.3		
2-b	75.0	Not Tacky	4.0	33.2		
2-с	80.9	Not Tacky	6.9	33.9		
3-a	69.5	Not tacky	3.9			·
3-b	76.6	Not Tacky		46.9	-31.5	-337

Gloss Measurement

- 1) Coat Leneta charts (Form N2C) with solution using a #8 Meyer rod
- 2) Allow chart to dry for 1 hour. Measure 60° gloss using a portable gloss meter at 3 points on the left 1/3 of the chart. Calculate the average of the 3 gloss values. Evaluate coating for tack, greasiness, fingerprint mark and if and how the coating rubs off the chart.
- 3) 4 hours after the drawdown was completed, measure 60° gloss using a portable gloss meter at 3 points on the middle 1/3 of the chart. Calculate the average of the 3 gloss values. Evaluate coating for tack, greasiness, fingerprint mark and if and how the coating rubs off the chart.
- 4) 24 hours after the drawdown was completed, measure 60° gloss using a portable gloss meter at 3 points on the right 1/3 of the chart. Calculate the average of the 3 gloss values. Evaluate coating for tack, greasiness, fingerprint mark and if and how the coating rubs off the chart.
- 5) Using the average gloss at the 3 different times, calculate the overall average.

Curl Retention Test Method

Materials

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- Prepared natural virgin brown hair tresses or oriental hair of 2 g, 25 cm.
 - Comb Trade Mark: Matador; reference 2618/6 GB.
 - Humidity chamber to regulate temperature and humidity during test.

Procedure for pre-treating the swatches (washing):

- 25 1) Wet 5 tresses for 30 sec with tap water at 37°C.
 - 2) Lather the 5 tresses for 30 sec. With 5 g of the 30% SLS solution (Empicol LX28/Albright & Wilson), stroking the tresses downward, ensure you repeat the same movement for all the hair washed. Leave the surfactant to act for 30 sec.
 - 3) Rinse the tresses for 1 minute with tap water at 37°C.
- 30 4) Remove the excess of water by running the tresses between the two fingers 3 times.
 - 5) Allow the tresses to dry overnight on a paper towel at room temperature.

Procedure for treating the swatches with the resin and curling:

Blank or negative control is the solvent used in the treatments.

- 35 1) Dip 1 tress at a time 3 times in 37 °C tap water and remove excess of water by stroking the tress between 2 fingers
 - 2) Lay the tress down on a clean support and apply 100 microliters of a 6% resins solution all along the using a calibrated micropipette
 - 3) Detangle the tress completely
- 40 4) Roll the tress on rod spiral curler,
 - 5) Leave the swatches to dry overnight in an oven at 40 °C.

Test - curl retention measurements:

- 1. Start the humidity chamber 2 hours before the test set at 70% humidity and 25 C.
- 2. Carefully remove the roller from hair by twisting it slightly, 10 minutes before the start of the test. Cut the ends of the tress in order to make it even (cut as few as possible).
- 3. Ensure each tress is correctly curled.
- 4. Hang the tresses in the humidity chamber: the bottom of the wax sealing on the tress should be on the line "0" of the millimeter paper sheet in the back of the chamber.
- 5. Measure hair tress length at predetermined intervals of time, the length is measured as the distance between the bottom of the wax sealing and the bottom of the tress be aware that the bottom of the tress is going down, so the view angle should also go down in order to avoid reading mistakes.
- 6. After the 5 hours, remove the tresses from the humidity chamber and measure the tress length at its maximum, by unrolling it completely. Calculate curl retention as described below.

Percent curl retention is calculated as follows:

max length - length at
$$T = x$$

% Curl Retention = ------ X 100
max length - length at $T = 0$

Instron Test for Hair Conditioning for Dry or Wet Hair

25 Materials

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- Testing was performed on slightly bleached European.
- Comb Trade Mark: Unbreakable or Ace Brand (Fine tooth spacing: 16 teeth/25 mm.)
 Wide tooth spacing: 11 teeth/25 mm.)

30 Hair swatches preparation and Baseline testing:

- 1) Weigh out between 2.35 and 2.50 g. hair
- 2) Rough up lower half of plastic tab with sandpaper
- 3) Cut $\sim 1/2$ " off root end of hair.
- 4) Put glue in middle of sanded end of tab and lay hair on the glue. Squeeze more glue on top of hair; then using another plastic tab press glue into hair while evenly spreading the hair and glue onto the tab keeping ~ 1/8 to 1/4 inch away from sides.
- 5) Cut tress to $6" \pm 0.25"$ from bottom of tab.
- 6) Let glue dry overnight.
- 7) Punch hole in middle of tab $\sim 1/4$ " from the top.
- 40 8) Wash all tresses according to the following procedure:
 - a) Wet hair tress under 40 degree C tap water for 15 seconds.
 - b) Using a pipette, apply 1.0 g of 9% Sodium Lauryl Sulfate (active) and stroke through tress for 30 seconds.
 - c) Rinse tress under the running water for 30 seconds.

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- d) Place tresses on paper towel covered tray and dry overnight.
- e) Comb through tress three times with narrow teeth.
- 9) Measure force it takes to comb the untreated tresses before treatment using an Instoni machine. Take force measurements on dry and wet hair as follows before treatment.

Ensure that the temperature and humidity inside the test room are appropriate for a testing (approximately 70 deg F and 30% RH).

- a) A 5 kilogram load cell is used for the testing
- b) Place 2 combs in the holders with wide-spaced teeth on the left and fine teeth on the right. Secure combs with holder screws so that approximately 1" of combs are sticking out of the front of the holder and over-lapping each other. For untreated tresses, use the same combs for combing all tresses; but for treated tresses, use different combs for every different treatment.
- c) Dry Combing Procedure: Detangle hair by combing tress 3 times, Retangle hair by swirling tress clockwise 3 times and counter clockwise 3 times, place tress on hanger and use Instron to comb through tresses.
- d) Wet Combing Procedure: Wet hair by dipping in distilled water and detangle hair by combing tress 3 times, retangle hair by dipping in distilled water three times, remove excess water by passing tress through index and middle fingers twice, place tress on hanger and use Instron to comb tress.

Procedure for treating and testing the swatches with resin solution:

- 1) Wet the tresses for 15 sec under tap water at 40°C.
- 2) Remove the excess water by pulling tress through index and middle fingers.
- 3) Lay the tresses on a clean support and apply 100 microliters of 6% active silicone product using a calibrated micropipette.
- 4) Allow solution to dry on hair 60 minutes for ethanol-based solutions, overnight for aqueous to test on dry hair.
- 5) Comb through tresses once before performing Instron study on both dry and wet hair see 9 from previous section.

Foundation Formulation

Pigment Premix:

50 wt% DC 245 Fluid

- 13.16 wt% Carde AS Titanium dioxide (caprylyl silane treated)
 - 11.41 wt% Carde AS Red Iron Oxide(caprylyl silane treated)
 - 18.26 wt% Carde AS-Yellow Iron Oxide(caprylyl silane treated)
 - 7.17 wt% Carde AS Black Iron Oxide(caprylyl silane treated)

10 Procedure:

- 1) Place DC 245 fluid in Waring Blender
- 2) Add titanium dioxide and mix by pressing the pulse button for 2 seconds for 15 seconds total.
- 3) Add red iron oxide and mix the same as titanium dioxide
- 15 4) Continue with the other pigments
 - 5) When all materials have been dispersed, mix on high and shred for 30 sec to grind the pigments
 - 6) Place premix into a round glass jar and place on pail roller for 6 hours.

20 Phase A

20.50 wt% Pigment Premix

7.50 wt% DC 5225C

8 wt% of a 50% resin solids in solvent

25 Phase B

54.80 wt % DI Water

1.0 wt% NaCl

0.20 wt% Polysorbate 20

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Procedure for Liquid Water in oil Foundation

- 1) Put pigment dispersion on roller for 1 hour.
- 2) Weigh out resin and solvent to make a 50% solids dilution. Use oven and wheel to mix
- 3) Combine ingredients in Phase A, mix until uniform using a dual blade, turbulent style mixing action.
 - 4) Combine ingredients in Phase B in separate beaker, mix until uniform using a magnetic stirrer
 - 5) Increase mixing speed of Phase A to 1376 rpm and very slowly add Phase B through an addition funnel. This addition should take 10 mins.

Continue mixing for an additional 10 min.

Foundation Durability Method: Gardner Abrasion Tester

- 1. Cut collagen into 3.5" x 3" pieces, place one on each of the 3" x 2.5" polycarbonate blocks and put in the humidity chamber overnight. This chamber must be at a constant 98% relative humidity level.
- 2. Remove collagen and block from chamber. Secure collagen to block with Scotch tape taking care not to place any tape on the top of the block's surface.
- 3. Add approximately 1 gram of foundation to the collagen, beading it across the top of the block. Using a #8 Meyer rod, coat the collagen with the foundation by placing the rod on the bead of foundation and spreading it downward to the bottom of the block. The final coating weight should be approximately 0.2 grams. This operation may need to be repeated to obtain the proper coating weight. Remove any material from the sides of the block.
- 4. Allow sample on collagen to dry. Drying times vary with different samples. Entire sample must be free from any wetness before testing. Measure color of sample on collagen for the initial baseline color using a spectrophometer or colorimeter. L*, a*, and b* designate the place of the colored object in a tri-dimensional space.
- 5. Place block with collagen face-up on the Gardner Abrasion Tester making sure that the block is in the tester. The soft side of Velcro is attached to the insult block to abrade or insult the foundation sample on the collagen. The insult block rubs back and forth across the foundation sample. One insult consists of one back and forth motion. Insult the foundation sample on the collagen 20 times. The machine can be stopped at certain intervals to measure the color.
- 6. After the foundation sample is insulted 20 times, the color is read as L*, a*, b* and the change in color, ΔΕ, is calculated (see equation below). The number of insults, coating weight, and repetitions can be changed to fit the needs of the material being tested. This is up to the discretion of the operator.
 - ΔL^* , Δa^* and Δb^* = value after abrasion value at initial baseline before abrasion.

 $\Delta E = (\Delta L^2 + \Delta a^{*2} + \Delta b^{*2})^{1/2}$

With larger ΔE 's, more foundation was removed and therefore the foundation is less durable.

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